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## (54) SUSTAINED RELEASE OF BIOLOGICALLY ACTIVE ORGANIC COMPOUNDS

We, LAPORTE INDUSTRIES LIMITED, a British Company, of (71)Hanover House, 14 Hanover Square, London, W.1, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following

This invention relates to sustained release of biologically active organic compounds, that is, organic compounds having utility by virtue of their biological

activity.

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Examples of biologically active organic compounds are for example, insecticides, fungicides, molluscicides, herbicides, acaricides, nematocides or plant growth regulants. Such compounds usually take effect on being absorbed in the form of a solution in water into a target organism. It may be desirable to maintain the presence of one or more biologically active compounds in absorbable form over an extended period of time for example, for protection of growing crops from attack by diseases and pests. This may be accomplished by utilising sustained release compositions containing a biologically active compound which may be released slowly as a result of the effect of environmental conditions such as the effect of temperature, or moisture on the composition.

Sustained release compositions are subject to exacting performance requirements. There is, generally, a fairly narrow acceptable range of release rates, 20 below which there is an insufficient concentration of the biologically active compound in absorbable form and above which there is an unacceptable degree of wastage. Preferably, therefore, the active compound is released from the sustained release composition at as uniform a rate as possible and at a rate which produces a 25 level of concentration of active compound as close as possible to the optimum.

The present invention provides a sustained release composition comprising an inorganic particulate substrate having a cation exchange capacity, cations ionically bonded to the substrate, the cations having the formula

R<sub>n</sub> X H<sub>m</sub><sup>+</sup>

where x represents a nitrogen, phosphorus, antimony or arsenic atom having a 30 valency of 4 or an oxygen, selenium, sulphur or tin atom having a valency of 3, R, represents one or more organic radicals at least one of which contains a carbon-carbon chain of at least 10 carbon atoms and  $H_m$  represents sufficient hydrogen atoms, if any, required to satisfy the valency of X, and a biologically active organic compound deposited onto the substrate. 35

Suitably the substrate has a structure based on silicon or phosphorus atoms and very suitably has a structure based on linked SiO<sub>4</sub> tetrahedra. Preferably the substrate comprises an anionic lattice comprising layers of linked SiO, tetrahedra possibly interleaved by layers of other minerals such as Mg(OH)<sub>2</sub> or Al(OH<sub>3</sub>), for example, a clay mineral. Suitable clay minerals may belong to any of the groups of minerals described in Kirk-Othmar, Encyclopaedia of Chemical Technology 2nd Edition, Volume 5, pages 541—557. Preferably the substrate is a member of the smectite group. Smectite clay minerals have structures based on that of the mineral

5	of Al <sup>3+</sup> ions sandwiched tetrahedra. The central A Al <sub>2</sub> (OH) <sub>6</sub> , in which 2 out o pseudo-hexagonal network octahedral (Al) and tetrah usually sodium or calcium about 0.66 additional mon	s of superimposed layers each of which contains a plane between two inward pointing sheets of linked SiO <sub>4</sub> Al <sup>3+</sup> section may be regarded as a layer of gibbsite. If 3 OH ions are replaced by apical oxygens of an Si <sub>4</sub> O <sub>10</sub> and the charge balance is upset by substitution in both ledral (Si) sites and is redressed by inter-layer cations, cations. A typical extent of such substitution requires ovalent cations per formula unit and these ions are in the smectite group includes the following groups of	5
	Montmorillonite	$Si_8Al_{3.34}Mg_{0.86}(\frac{1}{4}Ca,Na)_{0.86}$	
	Beidellite	$Si_{7.34}Al_{0.66}Al_{4}(\frac{1}{2}Ca,Na)_{0.66}$	
	Nontronite	$Si_{7.34}Al_{0.66}Fe_4^{3+}(\frac{1}{2}Ca,Na)_{0.66}$	-
- •	Saponite	$Si_{7.34}Al_{0.66}Mg_{e}(\frac{1}{2}Ca,Na)_{0.66}$	
15	Hectorite	$Si_{8}Mg_{5.34}Li_{0.88}(\frac{1}{2}Ca,Na)_{0.86}$	15
	Sauconite	$Si_{6.7}Al_{1.3}Zn_{4-5}(MgAl, Fe^{3+})_{2-1}(\frac{1}{2}Ca,Na)_{0.66}$	
20	and are particularly suitable exchangeable cations of the suitable clay mineral which is Wyoming bentonite. Alt by a process described in least or the suitable clay and the suitable clay mineral which is the suitable can be suitable can b	rillonite minerals have a high cation exchange capacity ble for use in the present invention. The majority of the ne mineral are preferably sodium cations. A particularly his, in its natural state, substantially in the sodium form, ernatively a synthetic clay mineral such as that produced British Patent No. 1054111 or British Patent No. 1213122 bly the exchangeable cations of the substrate may be	20
25	Preferably the cation and, particularly, at least 0 many montmorillonite cla- preference.	exchange capacity of the substrate is at least 0.5 m.eq/g 0.65 m.eq/g. Many smectite clay minerals and especially, y minerals have cation exchange capacities fulfilling this	25
30	The cations utilised in cations the element X has protonation, very suitably H <sub>m</sub> represents sufficient had valency of X. Particularly	n the invention are known as 'onium' cations. In such its highest possible valency. This may be achieved by under acid conditions, of the compound RXH <sub>m</sub> where sydrogen atoms, minus 1, to satisfy the highest possible suitable onium cations are based on nitrogen e.g. those	30
35	compounds. In the onium formula preferably contain suitable organic radicals hatoms are those derived	the corresponding amine, or quaternary ammonium n cations utilised in this invention R <sub>n</sub> in the general ns from 10 to 30 carbon atoms. Examples of particularly aving chains of average length of about 10 to 20 carbon from tallow, soya or coco oil. Nitrogen based onium ines bearing such radicals are available under the Trade	35
40	Name Duomeen. Preferab 3, or 4 organic radicals R, 10 carbon atoms and par The cation exchange	If y the onium cation is based on nitrogen and contains 2, 1 or 2 of which contain a carbon-carbon chain of at least ticularly preferably not more than 20 carbon atoms, capacity of the substrate is preferably saturated with the this the onium cation is preferably in excess over the	40
45	cation exchange capacity above 1.0 and up to 1.3 tir manufacture of any of the treated in the form of a dis	of the substrate. A suitable quantity of onlum cation is nest the cation exchange capacity of the substrate. In the compositions of this invention the substrate is preferably spersion in water at a concentration of, for example, from	45
50	efficient adduction that the shear assists dispersion are shear stirrer. Additionally may be included in the	ne onium cation. It is an important factor in achieving an e substrate be in a highly dispersed state. The presence of a suitable way of attaining this is by means of a high a dispersion agent, such as tetrasodium pyrophosphate, suspension. The quantity of dispersing agent may be	50
55	substrate. If any aggregate remove such aggregates b	1% to 5% and preferably from 1% to 4% by weight of the es of substrate remain undispersed it may be desirable to by, for example, centrifuging. The onium cation may be if the substrate or may itself be dispersed in water, at, for of from 1% to 6% by weight, and the two dispersions	55

mixed by slowly adding the suspension of the onium compound to the dispersion of the substrate. Preferably the dispersion of the substrate and the mixed dispersions are maintained under shear throughout, and for a sufficient time after the mixing has been completed to allow the cation exchange to go to completion. A suitable time is for up to 30 minutes after mixing has been completed. Preferably the temperature is maintained throughout at from 10°C to 90°C depending on the 5 5 thermal stability of the onium cation. The derivative resulting from the cation exchange may be filtered and washed free of inorganic cations from the substrate and anions associated with the onium cation. Again depending on the thermal stability of the onium cation it may be 10 10 necessary to control the temperature of drying carefully to avoid decomposition of The derivative resulting from the ion exchange of an onium cation onto an organic substrate and bearing a biologically active organic compound in intimate 15 association with the onium cations is capable of giving a controlled rate of release 15 of the biologically active organic compound. The inorganic substrate is inherently hydrophilic. The presence of the onium cation gives, to an extent, an organophilic character to the substrate which may facilitate, to an extent, the formation of the intimate association between the 20 substrate and the deposited active compound. It is possible also, by suitable 20 selection of the onium cation, so to reduce the hydrophilic character of the substrate as to assist in reducing the exposure of the active compound carried on it to moisture thereby reducing the tendency for the active compound to dissolve at an undesirably high rate. These effects make it possible to tailor the rate of release of active compound by simply selecting an onium cation having an organic chain or 25 25 chains of suitable configuration and chain length to give the desired balance of organophilic and hydrophilic properties. We also find that the particle size of the substrate has a marked effect on the rate of release of active compound. This parameter is particularly easy to control in the case of inorganic substrates such as clay minerals which are commonly available commercially in specified particle sizes or, if not, are easily sieved to the 30 30 desired size. If the onium cation/active compound combination selected has too high a rate of release it may be possible to reduce that rate by using a substrate consisting of somewhat larger particles than might otherwise have been selected. The presence of very small particles may give rise to practical difficulties in 35 35 distribution of the composition due to dusting but this may be alleviated by forming composite granules, or the like, containing an inert centre and bearing an outer coating comprising the composition of this invention and, if necessary, a suitable binder. In one aspect of this invention the biologically active organic compound may 40 40 be bound to the onium cation by a hydrolysable chemical linkage. In the presence of moisture the hydrolysable finkage tends to break and to release the active compound. This mechanism of release may have particular application where the biologically active compound is of a highly toxic nature since the formation of a chemical bond with one or more functional groups of the organic compound may 45 45 reduce its toxicity and that of the sustained release composition during storage and handling operations. It is important to ensure that the hydrolysable linkage breaks in preference to the linkage between the onium cation and the inorganic substrate since otherwise the released active compound will still have the onium cation attached to it with a resulting effect on the solubility of the released active 50 50 compound and also, probably on its activity. It is a very important feature of certain embodiments of this aspect of the invention that an ionic linkage between the siliceous inorganic substrate, and particularly a substrate comprising an anionic lattice comprising layers of linked SiO<sub>4</sub> tetrahedra, for example a clay mineral, and an onium cation can be particularly stable in the presence of water. As a result the 55 55 onium cations tend to remain ion exchanged on the inorganic substrate under conditions which result in the breakage of the hydrolysable linkage but which might be expected, also, to result in at least the partial breakdown of an ionic linkage. A hydrolysable linkage may be formed in a number of different reactions. The 60 60

hydrolysable salts:-

reaction of amines, or amides with carboxylic acids results in the formation of

## $R.COOH + H_2NOCR^1 - RCOO^{-1}H_3NOCR^1$

## $R.COOH + Cl^{-} + NH_{3}R \rightarrow RCOO^{-} + H_{3} NR^{3} + HCl$

5	employed in the practice Onium cations partic invention are those havi ammonium groups capa biologically active compo	of this inventi ularly suitable f ng, after ion e ble of reactin und, or free ac	on. or us xchai g wi id gr	e and are readily formed and may be the in the practice of this aspect of the tage to the substrate, free amine or the carboxylic acid groups on the the oups capable of reacting with amine	5
10	carboxylic acid group is Chemicals Limited, which crotonic acid. An examp	ompound converse Armeen Z (Transition is produced by the of a composite of a composite	ertibled the state of the state	ompound.  e to the onium form and having a Mark), available from Armour Hess reaction of primary coco amine and convertible to the onium form and Mark) which is also available from	10
15	Armour Hess and is an residues. The coco, tallow degree which makes thes It must be noted the	assymmetrical wor soya reside compounds sat while it is r	dian ues p suitab oossib	nine based on coco, tallow or soya provides organophilic properties to a ble for use in the present invention, ble, according to one aspect of the yeable chemical linkage between the	15
20	biologically active comp control over the rate of circumstances, there is in to be slightly higher than	ound and the of release, wh at least some in if the biologica	oniui ich p istan illy a	n cation to impart some degree of cossibility may be useful in some ces a tendency for the rate of release ctive compound is merely adsorbed, onto a substrate on which the onium	20
25	cation has been ion excha hydrolysable chemical l sustained release appears the presence of an intin	nged without the inkage therebe to be the pres nate association	nere l etwee ence n, ac	being any or any substantial degree of in. The major factor in achieving and nature of the onium cation and hievable by deposition as hereafter ompound and the onium cation.	25
30	Examples of biologi invention are listed in T compound which could be	cally active co able I which a be utilised to fo	mpo Iso i rm a	unds which may be utilised in this dentifies a functional group in each hydrolysable linkage with a suitable ivity associated with each compound	30
35		Н		Herbicide	35
		F		Fungicide	
		N	_	Nematocide	
		I	_	Insecticide	

- Plant Growth Regulant

PG

T	ABI	LE	1

		TABLE 1	
	Common Name	Chemical Name	Active as
	Aminotriazole	3-amino-1,2,4 triazole	н
v	Asulam	4-aminobenzenesulphonylcarbamate	Н
dnox	Dicloran	2,6 dichloro-4-nitroaniline	F
contain amine groups	Menazon	S-4,6-diamino-1,3,5-triazin-2-ylmethyl OO-dimethyl phosphorodithioate	I
ontain	Secbumeton	2-sec-butylamino-4-ethylamino-6- methoxy-1,3,5 triazine	н
	Terbutryne	2-tert-butylamino-4-ethylamino-6-methylthio- 1,3,5-triazine	н
a	Chloramben	3-amino-2,5 dichlorobenzoic acid .	н
	Dal apon	2,2-dichloropropionic acid	н
roups	2,4,5-T	2,4,5-trichlorophenoxyacetic acid	н
oid 8	Dichlosprop	2-(2,4-dichlorophenoxy)propionic acid	н
lic ad	Dicamba	3,6-dichloro-2-methoxybenzoic acid	н
boxy	2,4-D	2,4-dichlorophenoxyacetic acid	н
Contain carboxylic acid groups	Endothal	7-oxabicyclo(2,2,1)heptane-2,3-dicarboxylic acid	н
Cont	Clorfenac	2,3,6-trichlorophenylacetic acid	Н
		Naphthalene acetic acid	P, G
	Terbucarb	2,6-di-tert-butyl-p-tolylmethyl-carbamate	н
sdn	Monocrotophos	dimethyl cis-1-methyl-2-methyl carbamoylvinyl phosphate	I
e 810	Bufencarb	3-(1-methylbutyl)phenylmethyl-carbamate	I
amid	Carbaryl	l-naph thylmethyl carbamate	ī
tituted amide groups	Carbofuran	2,3-dihydro-2,2-dimethylbenzofuran-7-ylmethylcarbamate	I
ır subst	Dimethoate	O,O-dimethyl-S-methyl carbamoyl-methyl phosphorodithioate	Ī
ide o	Diuron	3-(3,4-dichlorophenyl)-1,1-dimethylurea	Н
Contain amide or substi	Methomyl	1-(methylthio) ethyliden eamino methyl carbamate	I
Con	Aldicarb	2-methyl-2-(methylthio)propylide- amino methylcarbamate	I, N
		Naphthalene acetamide	P, G

may be monitored colourimetrically.

In each Experiment a cylinder of Perspex (Trade Mark) 70 mm in diameter and fitted with a closure in the lower end, the closure being fitted with a drain tube

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layers of material: Chelford Medium Sand — I cm depth approx. Glasswool — 1.5 cm depth approx.
Whatman No. 3 (Trade Mark) Filter Paper
Chelford Medium Sand — 10 cm approx. 2. 3. 5 5 Sustained release composition mixed with 50 ml Chelford Medium Sand. 5. Chelford Medium Sand — 1 cm depth approx. 6. Whatman No. 3 Filter Paper The sustained release composition used in each Experiment was produced as 10 10 follows: EXPERIMENT I 50 g of calcium montmorillonite granules available from Laporte Industries Limited as SYK 22/44 Granules (the figures refer to the British Standard sieve mesh size range of the granules which corresponds to a particle size; i.e. BSS; of from 355 15 15 to 760 microns) was mixed with 3 g of methyl orange pigment which had previously been dissolved in 250 g of distilled water. The resulting slurry without first removing excess moisture was dried in an oven. The oven dried granules were found to have a skin of dried methyl orange which was removed by passing the granules through a 22 BSS sieve. The final loading of methyl orange on the granules 20 20 was 24.75 mg/g. No onium cation was included. **EXPERIMENT 2** The method used was the same as for Experiment 1 but using a sieve fraction of from 85 to 100 mesh BSS obtained by grinding and sieving the granules used in Experiment 1. This corresponds to a particle size range of from 150 to 180 microns. 25 25 The final loading of methyl orange on the granules was 16.50 mg/g. No onium cation was included. EXPERIMENT 3 22.75 g of sodium bentonite belonging to the montmorillonite group of minerals available under the Trade Mark Volclay and having a cation exchange 30 30 capacity of 0.79 milliequivalents/g (me/g) and a particle size range of from 32-53 microns and being equivalent to a dried weight of volclay of 20.0 g was dispersed in 400 mls distilled water and the dispersion was heated to 80°C. 5.76 g of an unsymmetrical diamine based on tallow oil available under the Trade Mark Duomeen was dispersed in 16 mls of 1 N HCl (sufficient in theory to protonate one 35 35 of the two amine groups of the Duomeen to give an onium cation) and 200 mls distilled water and the dispersion was heated to 80°C. The two dispersions were mixed and stirred for 15 minutes, to allow the Duomeen, preferentially protonated at the secondary amine group, to ion exchange with sodium ions on the Volclay. The resulting adduct was filtered and washed free of chloride ions. 2.5 g methyl 40 40 orange was dissolved in 250 g distilled water and mixed with the adduct which was filtered to remove excess moisture and dried in an oven. The final loading of methyl orange on the Volclay was 58.9 mg/g. Since the free amine groups of the Duomeen were unprotonated, except to such extent as may occur naturally in an aqueous medium, it is thought that no chemical bond, or substantially none, was formed 45 45 between the methyl orange and the onium cation. **EXPERIMENT 4** The method was the same as that used for Experiment 3 except that the Volclay had a particle size range of below 32 microns. The final loading of methyl 50 orange was again 58.9 mg/g. 50 EXPERIMENT 5 The method used was the same as for Experiment 4 except that the Duomeen was replaced by 10.78 g of dimethyl dioctadecyl ammonium chloride (available under the Trade Mark Arquad 2HT) and no acid was used since the Arquad 2HT, being a quaternary ammonium compound, is already suitable for adduction onto 55 55 the Volclay. The final loading of methyl orange on the Volclay was 56.4 mg/g. No chemical bond could have formed between the methyl orange and the alkyl chains

of the onium cation.

having a tap, was positioned vertically and loaded successively with the following

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**EXPERIMENT 6** 

The method used was the same as for Experiment 3 except that 32 mls of 1N HCl wa used so as to protonate both amine groups of the Duomeen. The final loading of methyl orange on the Volclay was 147.69 mg/g. It is thought that a substantial proportion of the methyl orange would have been bonded to free protonated primary amine groups on the onium cation by salt formation with the —SO<sub>3</sub> residues on the methyl orange.

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EXPERIMENT 7

The method used was the same as for Experiment 6 except that the Volclay had a particle size range of below 32 microns. The final loading of methyl orange was again 147.69 mg/g. Again it is thought that a substantial proportion of the methyl orange would have been chemically bonded to the onium cation.

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The dosage of methyl orange included in layer number 5 of the column in each of Experiments 1 to 5 was 2.31 mg and in each of Experiments 6 and 7 was 2.6 mg and the appropriate quantity of sustained release composition to attain this dosage of methyl orange was used.

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In the course of the Experiments a series of aliquots of 400 ml of distilled water were added to each column and the outlet tube was choked by means of the tap to give a dropwise flow from the column at a rate of 400 ml over 12 hours. Samples of the eluant were examined colourimetrically and the quantity of methyl orange eluted with each aliquot of distilled water was determined. The results expressed as a cumulative % methyl orange remaining are summarised in Table 2 over a series of 12 aliquots identified in column 1.

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TABLE 2

Experiment			Wt.	% Remaini	ng		
Aliquots	1	2	3	4	5	6	7
1	76.84	62.99	97.38	95.58	75.58	95.85	82.19
2 .	48.44	41.35	93.46	85.23	51.38	84.47	53.04
3.	41.95	30.57	89.13	75.10	40.47	71.43	36.92
4	39.28	24.25	84.50	66.88	34.24	60.58	27.31
5	37.63	19.13	79.38	59.39	29.87	50.20	19.09
6	35.93	14.50	74.06	52.38	27.00	41.28	14.55
7	35.79	12.94	69.84	48.18	26.06	39.02	13.63
8	35.79	10.56	66.38	45.06	24.94	35.75	12.63
9	34.84	6.97	61.01	40.82	22.78	30.60	10.82
10	34.84	5.17	56.81	37.14	21.13	27.83	9.90
11	34.84	3.05	53.48	34.93	19.49	25.45	9.48
12	_	2.06	50.19	33.11	18.93	23.84	9.06
Remarks	No onium cation	No onium cation	Little or no chemical bonding	Little or no chemical bonding	No chemical bonding	Substantial Chemical Bonding	Substantial Chemical Bonding

5	The following Examples illustrate the invention. Examples 1 and 2 are according to the invention and Example 3 is not according to the invention but is for comparative purposes.  In each Example a leaching column 7 cm in diameter and loaded as described below was used.				
		Depth of Layers — listed from top of column	5		
	2.5 cm	0.7 diameter glass beads.			
		Whatman No. 541 filter paper.			
	10 cm	Chelford Medium Grade Sand.			
10	1.8 cm	Mixture of sustained release composition and sand.	10		
	10 cm	Chelford Medium Grade Sand.			
	8.8	Whatman No. 541 Filter Paper.			
15	capacity of of mono p sufficient ac 2,4-dichloro by weight a was 36-44 mineral was of Example	ading of sustained release composition was 0.24 g. In Examples 1 and 2 and release composition consisted of Volclay, having a cation exchange approximately 0.8 me/g, which had been ion exchanged with 0.75 me/g protonated Duomeen and which had thereafter been treated with a tid to protonate the free amine group of the Duomeen and treated with sphenoxyacetic acid (2.4-D). The Volclay carried 12.27% of the 2.4-D gainst a theoretical maximum of 11.5%. In Example 1 the clay mineral BSS mesh size (420—345 microns diameter). In Example 2 the clay 60—100 BSS mesh size (250—150 microns). Example 3 was a repetition 2 except that no Duomeen was used and the clay mineral was not acid	15		
25	flow slowly further aliqu four eluates	course of the Examples a 400 ml aliquot of distilled water was allowed to through the column in the course of a day and this was repeated using a not of 400 ml distilled water on each of 3 subsequent days. Each of the was examined as follows. A 25 ml sample of the eluate was extracted 0 ml CCl <sub>4</sub> . The CCl <sub>4</sub> was evaporated to 0.1 ml on a steam bath and 5.0	25		
30	evaporated sample was cooled and	to dryness and 5 ml of concentrated sulphuric acid was added. The then heated at 130°—135°C for 20 minutes. The sample was then made up to 50 ml with distilled water. If 2.4-D is present the sample is	30		
35	an estimate entire eluate	red. Each sample is examined colourimetrically using a Eel eter — No. 606 filter — at 565 mm in an appropriate size cell to enable of the quantity of 2.4-D present in the sample and, thereby, in the e, to be made by reference to a calibration curve which showed near er the range 0—25 ppm 2,4-D.	35		

linearity over the range 0—25 ppm 2,4-D.

The results, expressed as a cumulative weight % of the 2,4-D originally present in the sustained release composition are shown in Table 2.

TABLE 2

		Example No.			
Eluate No.	1	2	3		
1	17	35.5	33.5		
2	51.5	76	96.5		
3	62.5	83	97		
4	69.5	86.5	97.5		

	WHAT WE CLAIM IS:—	
	<ol> <li>A sustained release composition comprising an inorganic particulate substrate having a cation exchange capacity, cations ionically bonded to the substrate, the cations having the formula</li> </ol>	
5	$R_n \times H_m^+$	5
10	where X represents a nitrogen, phosphorus, antimony or arsenic atom having a valency of 4 or an oxygen, selenium, sulphur or tin atom having a valency of 3, R <sub>n</sub> represents one or more organic radicals at least one of which contains a carbon-carbon chain of at least 10 carbon atoms and H <sub>m</sub> represents sufficient hydrogen atoms, if any, required to satisfy the valency of X, and a biologically active organic compound deposited onto the substrate.  2. A composition as claimed in claim 1 wherein the substrate is a clay mineral.	10
15	3. A composition as claimed in claim 2 wherein the substrate is a clay mineral of the smectite group.  4. A composition as claimed in claim 3 wherein the substrate is a clay mineral of the montmorillonite group.	15
20	<ul> <li>5. A composition as claimed in any preceding claim wherein the substrate has a cation exchange capacity of at least 0.5 m.eq/g.</li> <li>6. A composition as claimed in any preceding claim wherein, in the formula. X represents a nitrogen atom.</li> <li>7. A composition as claimed in claim 6 wherein, in the formula, n is equal to 2,</li> </ul>	20
25	3 or 4 and one or two of the organic radicals represented by R <sub>n</sub> contains a carbon-carbon chain of at least 10 carbon atoms.  8. A composition as claimed in claim 7 wherein in the formula each organic radical R contains not more than 30 carbon atoms.  9. A composition as claimed in any preceding claim wherein the biologically active organic compound is bound to the cations by a hydrolysable chemical	25
30	linkage.  10. A composition as claimed in any preceding claim wherein the cation exchange capacity of the substrate is saturated by the cations.  11. A composition as claimed in any preceding claim wherein the biologically active compound is an insecticide.	30
35	12. A compound as claimed in claim 11 wherein the insecticide is S-4,6-diamino-1,3,5-triazin-2-ylmethyl OO-dimethyl phosphorodithioate.  13. A composition as claimed in claim 11 wherein the insecticide is dimethyl cis-1-methyl-2-methyl carbamovlvinyl phosphate.  14. A composition as claimed in claim 11 wherein the insecticide is 3-(1-	35
40	methylbutyl) phenylmethylcarbamate.  15. A composition as claimed in claim 11 wherein the insecticide is 1-naphthylmethylcarbamate.  16. A composition as claimed in claim 11 wherein the insecticide is 2,3-dihydro-2,2-dimethylbenzofuran-7-ylmethylcarbamate.	40
45	17. A composition as claimed in claim 11 wherein the insecticide is O,O-dimethyl-S-methylcarbamoylmethyl phosphorodithioate. 18. A composition as claimed in claim 11 wherein the insecticide is 1-(methylthio)ethylideneamino methylcarbamate. 19. A composition as claimed in claim 11 wherein the insecticide is 2-methyl-2-	45
50	(methylthio) propylideamino methylcarbamate.  20. A composition as claimed in any one of claims 1 to 10 wherein the biologically active compound is a herbicide.  21. A composition as claimed in claim 20 wherein the herbicide is 3-amino-124 triazole	50
55	22. A composition as claimed in claim 20 wherein the herbicide is 4-aminobenzenesulphonylcarbamate.  23. A composition as claimed in claim 20 wherein the herbicide is 2-secbutylamino-4-ethylamino-6-methoxy-1,3,5 triazine.  24. A composition as claimed in claim 20 wherein the herbicide is 2-tert-	55
60	butylamino-4-ethylamino-6-methylthio-1,3,5-triazine.  25. A composition as claimed in claim 20 wherein the herbicide is 3-amino-2,5 dichlorobenzoic acid.  26. A composition as claimed in claim 20 wherein the herbicide is 2,2-	60
	dichloropropionic acid	

11	1,363.362	11
	27. A composition as claimed in claim 20 wherein the herbicide is 2,4.5-	-
	trichlorophenoxyacetic acid.  28. A composition as claimed in claim 20 wherein the herbicide is 2-(2,4-	
	dichlorophenoxy) propionic acid.	
5	29. A composition as claimed in claim 20 wherein the herbicide is 3.6-dichloro-	5
	2-methoxybenzoic acid.  30. A composition as claimed in claim 20 wherein the herbicide is 2.4-	
	dichlorophenoxyacetic acid.	
	31. A composition as claimed in claim 20 wherein the herbicide is 7-	
10	oxabicyclo(2,2,1)heptane-2,3-dicarboxylic acid.  32. A composition as claimed in claim 20 wherein the herbicide is 2,3,6-	10
	trichlorphenylacetic acid.	
	33. A composition as claimed in claim 20 wherein the herbicide is 2.6-di-tert-	
	butyl-p-tolymethyl-carbamate.  34. A composition as claimed in claim 20 wherein the herbicide is 3-(3,4-	15
15	dichlorophenyl)-1,1,-dimethylurea.	13
	35. A composition as claimed in any one of claims 1 to 10 wherein the	
	hiologically active compound is a fungicide.	
20	36. A composition as claimed in claim 35 wherein the fungicide is 2,6 dichloro- 4-nitroaniline.	20
24	37. A composition as claimed in any one of claims 1 to 10 wherein the	
	hiologically active compound is a nematocide.	
	38. A composition as claimed in claim 37 wherein the nematocide is 2-methyl-2-(methylthio)propylideamino methylcarbamate.	
25	39. A composition as claimed in any one of claims I to 10 wherein the	25
	hiologically active compound is a plant growth regulant.	
	40. A composition as claimed in claim 39 wherein the plant growth regulant is naphthalene acetic acid.	
	41. A composition as claimed in claim 39 wherein the plant growth regulant is	
30	nanhthalene acetamide.	30
	42. A composition as claimed in any one of claims 1 to 10 wherein the biologically active compound is an organometallic compound.	
	43. A composition as claimed in claim 42 wherein the biologically active	
25	compound is an organotin compound.	35
35	44. A composition as claimed in claim 43 wherein the biologically active compound is tributyl tin oxide.	33
	45. A composition as claimed in any preceding claim wherein the biologically	
	active compound is in the composition in from 10 to 250 mg per gram of substrate,	
40	the substrate being weighed when dry and without inclusion of the onium compound.	40
	46. A composition as claimed in claim 1 and substantially as described herein.	
	47. A process for treating a biological entity with an organic compound having	
	biological activity towards said entity comprising exposing the habitat of the biological entity to contact with water which has acquired a content of said organic	
45	compound as a result of contact with a composition as claimed in any preceding	45
	claim.	
	48. A process as claimed in claim 47 wherein the organic compound is a herbicide and the composition is applied to the vicinity of a plant towards which	
	the herbicide has no activity to protect the said plant from competing growth.	
50	49. A process as claimed in claim 48 wherein the plant to be protected is a tree	50
	sapling. 50. A process as claimed in claim 47 wherein the composition is included in a	
	surface coating medium and wherein the medium is applied to a surface subject to	
	contamination by the growth thereon of biological entities.	55
55	51. A process as claimed in claim 47 and substantially as described herein.	<b>33</b>

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